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## Drone data reveal heterogeneity in tundra greenness and phenology not captured by satellites

Jakob J Assmann<sup>1,2</sup> , Isla H Myers-Smith<sup>1</sup> , Jeffrey T Kerby<sup>3</sup> , Andrew M Cunliffe<sup>4</sup> and Gergana N Daskalova<sup>1</sup>

<sup>1</sup> School of GeoSciences, University of Edinburgh, Edinburgh, United Kingdom

<sup>2</sup> Department of Biology, Aarhus University, Aarhus, Denmark

<sup>3</sup> Aarhus Institute of Advanced Studies, Aarhus University, Aarhus, Denmark

<sup>4</sup> Department of Geography, University of Exeter, Exeter, United Kingdom

E-mail: [j.assmann@bio.au.dk](mailto:j.assmann@bio.au.dk)

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Supplementary material for this article is available [online](#)

### Abstract

Data across scales are required to monitor ecosystem responses to rapid warming in the Arctic and to interpret tundra greening trends. Here, we tested the correspondence among satellite- and drone-derived seasonal change in tundra greenness to identify optimal spatial scales for vegetation monitoring on Qikiqtaruk—Herschel Island in the Yukon Territory, Canada. We combined time-series of the Normalised Difference Vegetation Index (NDVI) from multispectral drone imagery and satellite data (Sentinel-2, Landsat 8 and MODIS) with ground-based observations for two growing seasons (2016 and 2017). We found high cross-season correspondence in plot mean greenness (drone-satellite Spearman's  $\rho$  0.67–0.87) and pixel-by-pixel greenness (drone-satellite  $R^2$  0.58–0.69) for eight one-hectare plots, with drones capturing lower NDVI values relative to the satellites. We identified a plateau in the spatial variation of tundra greenness at distances of around half a metre in the plots, suggesting that these grain sizes are optimal for monitoring such variation in the two most common vegetation types on the island. We further observed a notable loss of seasonal variation in the spatial heterogeneity of landscape greenness (46.2%–63.9%) when aggregating from ultra-fine-grain drone pixels (approx. 0.05 m) to the size of medium-grain satellite pixels (10–30 m). Finally, seasonal changes in drone-derived greenness were highly correlated with measurements of leaf-growth in the ground-validation plots (mean Spearman's  $\rho$  0.70). These findings indicate that multispectral drone measurements can capture temporal plant growth dynamics across tundra landscapes. Overall, our results demonstrate that novel technologies such as drone platforms and compact multispectral sensors allow us to study ecological systems at previously inaccessible scales and fill gaps in our understanding of tundra ecosystem processes. Capturing fine-scale variation across tundra landscapes will improve predictions of the ecological impacts and climate feedbacks of environmental change in the Arctic.

### 1. Introduction

Identifying the scales at which ecological processes operate is a fundamental, yet often neglected element of ecological research [1–3]. Cross-scale ecological information can inform our understanding of the causes and consequences of global change [2]. In tundra ecosystems, vegetation responses triggered by rapid Arctic warming could influence ecosystem

functions through altered carbon and nutrient cycles with potential feedbacks to the global climate system [4–8]. Yet, challenging logistics have limited the extent of field-based observations in Arctic ecosystems [9–11]. The grain sizes of global-extent satellite products (tens of meters to kilometres) are too coarse to capture the fine-scale dynamics of tundra plants [12–14] and to link vegetation change to key ecosystem functions [13]. Thus, by bridging this ‘scale-gap’,

we can transform our understanding of pan-Arctic tundra vegetation change and associated global-scale climate feedbacks.

### 1.1. Satellites show greening of the tundra

Satellite observations indicate a ‘greening’ of tundra ecosystems [13, 15–20] and shifts in growing season phenology over recent decades [21–24]. Observations of increasing tundra greenness are often reported from surface-reflectance-derived Normalised Difference Vegetation Index (NDVI) [16, 18, 25, 26]. Satellite-observed tundra greening has occurred concurrently with ground-based observations of vegetation change in Arctic ecosystems [27] including increased shrub cover [28–31] and taller community-level plant height [32], as well as earlier leaf emergence and flowering at some [33–36], but not all tundra sites [36–39]. However, mismatches between ground and satellite-based observations suggest the potential for an observational scale gap [13].

### 1.2. Arctic vegetation change and phenology have been linked to warming

Satellite-observed Arctic greening trends have been linked directly to warming air temperatures [19, 20, 40–46] and indirectly to sea-ice declines [17, 47–51]. Ground-based observations of tundra vegetation change correspond with warming [27, 32, 52], but do not always co-occur with satellite greening trends in the regions around the ecological monitoring sites [13, 53]. While satellite-based phenology observations from the Arctic have been mainly linked to temperature [22, 54, 55], *in situ* phenology in the tundra has been shown to be influenced by a suite of interacting factors rarely tested in satellite-based analysis of Arctic phenology. These factors include, but are not limited to: snowmelt, temperature, day length, and the proximal influences of sea-ice on localised climate affect [34–36, 38, 56, 57]. Thus, ecological studies indicate greater complexity of drivers than analyses of satellite-derived greening trends to date.

### 1.3. Inconsistencies amongst satellite platforms and heterogeneous greening trends

Greening trends and phenology measures derived from different satellite platforms do not always correspond with each other [13, 18]. Additionally, satellite-derived greening trends vary at global [18], continental [42, 58–60] and regional scales [46–48, 61–64]. Many areas of the Arctic show no trends in NDVI, with only around 20% of the Arctic spectrally greening and around 1%–4% of the Arctic spectrally browning [13, 62, 65, 66]. Recent analyses suggest a slowdown of the Arctic-wide spectral greening trend over the past decade [43, 67]. Furthermore, despite NDVI being related to the photosynthetically active biomass in the tundra [14, 68–70], geophysical, environmental and ecological factors,

such as low solar angle, atmospheric effects (including cloud and fog), snow cover, soil moisture and standing water, in addition to the non-linearity of NDVI-biomass relationships, complicate the interpretation of satellite-derived NDVI time-series at high latitudes [13, 71]. The growing complexity highlighted in Arctic greening trends has led to repeated calls for ground validation of satellite observations [11, 18, 59, 60, 66, 72, 73].

### 1.4. The scale discrepancy problem in Arctic greening

A major problem in linking satellite-derived trends of spectral greenness and phenology to *in situ* observations of ecological processes in the tundra is the discrepancy in observational scales [13, 29, 61, 72, 74]. Satellite datasets with long-term records are limited by their moderate- to coarse-grain sizes, ranging from 30 m (Landsat) to 250 m (MODIS) and 8 km (AVHRR-GIMMS3g). *In situ* ecological monitoring in the Arctic is logistically challenging and therefore restricted in extent to a limited number of sites and often metre-squared plots [10, 75]. Only a few studies have linked on-the-ground vegetation or phenology change to satellite trends in NDVI in Arctic tundra [13, 14, 47, 48, 53, 76–78]. However, drones equipped with compact sensors now allow for the collection of ultra-fine-grain multispectral imagery at landscape extents that can potentially bridge the scale-gap between satellite and ground-based observations [14, 79–82].

### 1.5. Novel drone data to study variation in greenness

Here, we set out to test whether drones can be used to identify the key ecological scales for studying tundra greenness on Qikiqtaruk in the Canadian Arctic by bridging the scale gap between satellite and *in situ* data. First, we tested whether satellite- and drone-derived measures of mean landscape-scale greenness (NDVI) agree across two growing seasons while controlling for the potentially confounding effects of topography and land cover. Second, we identified the key spatial scales for ecological variation in landscape greenness within the two most common vegetation types at our study site using variograms. Third, we tested how the magnitude of seasonal variation in tundra greenness scales across grain sizes from fine-resolution drone imagery to medium-grain satellite imagery. Finally, we assessed whether drone-derived NDVI corresponds with on-the-ground measures of within growing season change in plant growth based on methods frequently used by long-term, field-based monitoring networks. Thus, in our analysis, we validated satellite-derived landscape estimates of vegetation greenness with ultra-fine-grain drone data and described spatial and temporal variation in tundra productivity at landscape extents (1–100 ha) with grain sizes that were previously not accessible.

## 2. Methods

### 2.1. Site description: Qikiqtaruk—Herschel Island

Qikiqtaruk (69.57 N, 138.91 W) is located in the Beaufort Sea along the coastline of the North Slope of the Yukon Territory, Canada. The vegetation is the moist acidic shrub tundra [83] characteristically found in the Western Arctic regions of North America, which has experienced strong spectral greening in recent decades [13]. The two most common plant communities on the island are the tussock sedge ('Herschel') and *Dryas-vetch* ('Komakuk') vegetation types [84, 85]. The tussock sedge vegetation is dominated by the name-giving tussock sedge *Eriophorum vaginatum* L. with varying cover of *Salix pulchra* Cham. The top-soils of the island are underlain by ice-rich permafrost and undergo frequent disturbance [85]. The *Dryas-vetch* vegetation is particularly found on ground disturbed by soil creep and is characterised by the near ubiquitous presence of *Dryas integrifolia* Vahl., the willow *Salix arctica* Phall., various grass species including *Arctagrostis latifolia*. (R.Br.) Griseb. and forb species [86]. The relative abundances of these species are shown in (figure S1, which is available online at (<https://stacks.iop.org/ERL/15/125002/mmedia>)). Though the two vegetation types are specific to the region, these plant communities would group with tundra types S1, W2 and G3/4 of the Circumpolar Arctic Vegetation Map [87].

We established four study areas on the east end of the island, each with two co-located one-hectare plots in the two key vegetation types (figure 1, table S1). We selected plots with homogenous terrain and land cover to represent the two key vegetation types and to control for the potentially confounding effects of terrain and cover heterogeneity. The island harbours small herds of caribou (100s of individuals) and muskox (3–35 individuals in recent years) of fluctuating size, as well as cyclic populations of voles and lemmings [88]. We estimate the overall impact of herbivory on the vegetation in our study plots to be low particularly in 2016 and 2017 when there were few muskox on the island.

### 2.2. Multispectral drone time-series

We analysed a total of 62 drone surveys from 21 dates; see table S2 for a breakdown by one-hectare monitoring plots. We collected multispectral drone imagery using Parrot Sequoia (Paris, France) compact multispectral sensors mounted on multi-rotor drone platforms in June to August in 2016 and 2017. We used three different drone platforms: a Tarot 680 Pro hexacopter with camera sensor stabilisation in 2016, and a 3DR Iris + and a DJI Phantom 4 Pro without sensor stabilisation in 2017. Surveys were flown using parallel flight lines (a lawn-mower flight pattern) at an altitude of ca. 50 m, giving ground-sampling distances of 0.04 m to 0.06 m. Images were acquired with 75%

front- and side-lap as close as possible to solar noon (mean absolute difference to solar noon 2.16 h, maximum 6–7 h). See table S2 and the methods section of the supplementary materials for further details on the drone surveys, including additional information on radiometric calibration, as well as temporal and spatial coverage. Drone data and code to carry out all analyses are available via a GitHub repository [91].

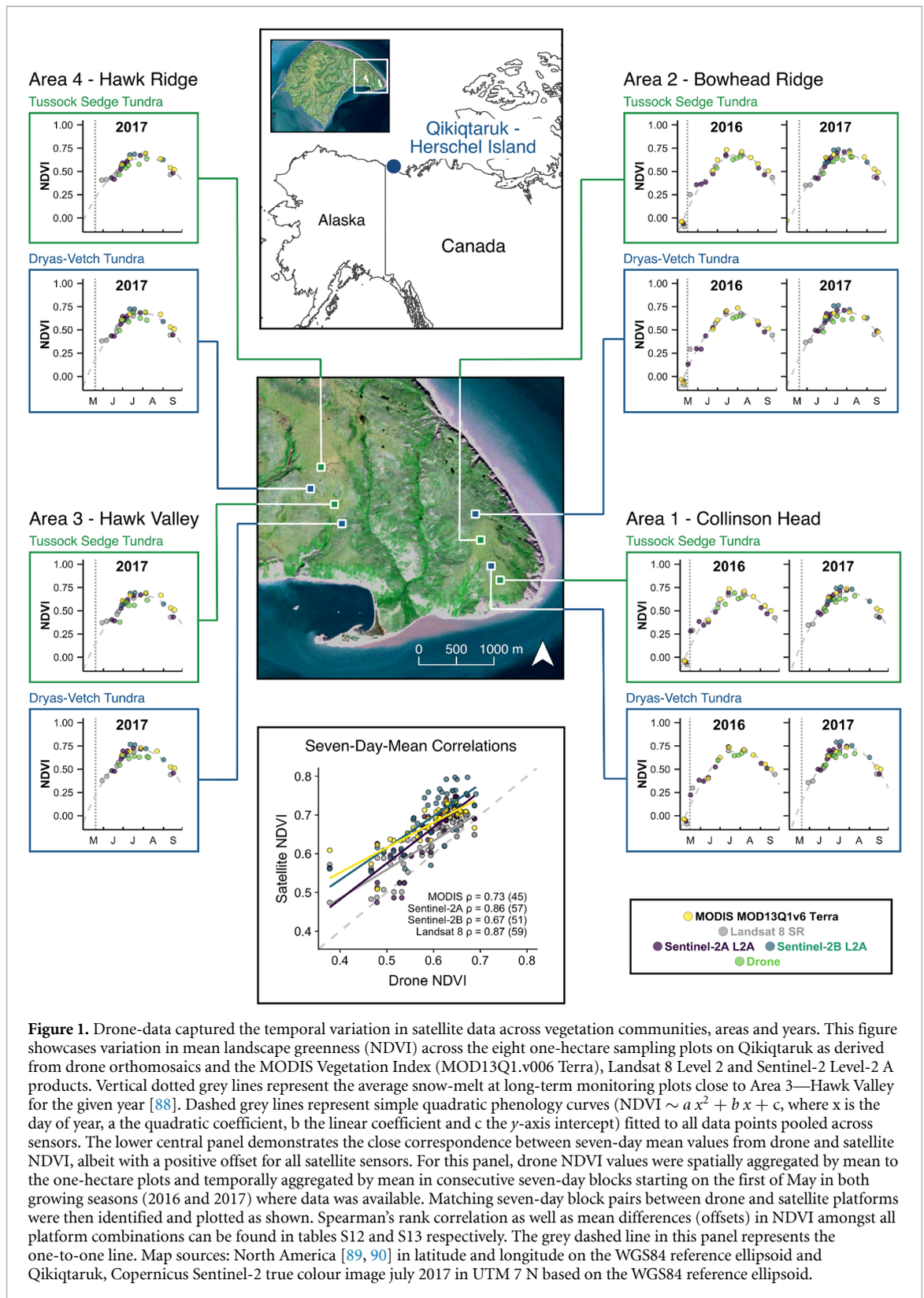
We processed the Sequoia imagery using Pix4D Mapper v4.0.21 (Lausanne, Switzerland) with the *agMultispectral* template and the 'merge map tiles' option set to true to generate co-registered single-band surface reflectance maps. Radiometric calibration was carried out in Pix4D Mapper using pre- or post-flight imagery of calibrated reflectance panels; in 2016 we used a MicaSense (Seattle, USA) panel and in 2017 a SphereOptics (Herrsching, Germany) Zenith Lite panel. We measured panel reflectance pre- and post-season and used the mean values for radiometric calibration. We also calibrated for sensor properties and sun irradiance measured by the incident light sensor. We used four to six ground control points per survey that were precisely geolocated with a GNSS system to spatially constrain the reconstructions in Pix4D Mapper with an estimated accuracy of 1–2 pixels between bands and 2–6 pixels between surveys [81]. We calculated the Sequoia NDVI as the normalised difference between the near-infrared (770 nm–810 nm) and red (640 nm–680 nm) bands of sensor.

### 2.3. Satellite time-series

Satellite time-series were obtained from three different satellite sensors: 1) the Moderate Resolution Imaging Spectroradiometer (MODIS) on the USGS Terra satellite, 2) the Multispectral Instrument (MSI) on Sentinel-2 A & B and 3) the Operational Land Imager (OLI) on Landsat 8.

We obtained MODIS NDVI values for the time period from the 1st May to the 30th of September in 2016 and 2017 for all 250 m MODIS pixels that contained the survey plots. NDVI values were retrieved from the 16-day MOD13Q1 v6 Terra product [92] using the Google Earth Engine [93]. We discarded all values with a 'Summary QA' score of –1 (no data) or 3 (cloudy). table S3 lists the resulting MODIS-pixel-date pairs. The MODIS NDVI is calculated as the normalised difference between bands 1 (841 nm–876 nm) and band 2 (620 nm–670 nm).

For the Sentinel-2 time-series, we gathered all Sentinel-2 MSI L1C scenes containing the tile covering Qikiqtaruk (T07WET) that were available on the Copernicus Open Access Hub (<https://scihub.copernicus.eu/>) for the same time period as the MODIS pixels. We processed all scenes to L2A using Sen2Cor 2.4.0 [94], retained all bands with 10 m resolution (2, 3, 4 and 8), applied the cloud mask and generated a true-colour image. We inspected all scenes visually and discarded all imagery with cloud contamination over the study area (78% of



**Figure 1.** Drone-data captured the temporal variation in satellite data across vegetation communities, areas and years. This figure showcases variation in mean landscape greenness (NDVI) across the eight one-hectare sampling plots on Qikiqtaruk as derived from drone orthomosaics and the MODIS Vegetation Index (MOD13Q1.v006 Terra), Landsat 8 Level 2 and Sentinel-2 Level-2 A products. Vertical dotted grey lines represent the average snow-melt at long-term monitoring plots close to Area 3—Hawk Valley for the given year [88]. Dashed grey lines represent simple quadratic phenology curves ( $NDVI \sim ax^2 + bx + c$ , where  $x$  is the day of year,  $a$  the quadratic coefficient,  $b$  the linear coefficient and  $c$  the  $y$ -axis intercept) fitted to all data points pooled across sensors. The lower central panel demonstrates the close correspondence between seven-day mean values from drone and satellite NDVI, albeit with a positive offset for all satellite sensors. For this panel, drone NDVI values were spatially aggregated by mean to the one-hectare plots and temporally aggregated by mean in consecutive seven-day blocks starting on the first of May in both growing seasons (2016 and 2017) where data was available. Matching seven-day block pairs between drone and satellite platforms were then identified and plotted as shown. Spearman's rank correlation as well as mean differences (offsets) in NDVI amongst all platform combinations can be found in tables S12 and S13 respectively. The grey dashed line in this panel represents the one-to-one line. Map sources: North America [89, 90] in latitude and longitude on the WGS84 reference ellipsoid and Qikiqtaruk, Copernicus Sentinel-2 true colour image July 2017 in UTM 7 N based on the WGS84 reference ellipsoid.

scenes for 2016 and 74% of scenes for 2017). The resulting set contained nine cloud-free Sentinel-2 L2A scenes of the study area from 2016 and 15 scenes from 2017 (table S4). Finally, the Sentinel NDVI was calculated as the normalised difference between band 8 (784.5 nm–899.5 nm) and band 4 (650 nm–680 nm).

Landsat 8 OLI Level-2 (surface reflectance) time-series were obtained using the USGS EarthExplorer website (<https://earthexplorer.usgs.gov/>) by querying the search engine for all scenes that covered the study site during the same time-period as the MODIS pixels ( $n = 94$ ). The automatically generated cloud masks were of insufficient quality, so we manually inspected

all scenes and retained only the scenes cloud-free over the study site ( $n = 7$  for 2016,  $n = 8$  for 2017, table S5). The Landsat 8 NDVI was then calculated as the normalised difference between band 5 (845–885 nm) and band 4 (630–680 nm). The study plots were not designed with a Landsat 8 analysis in mind and did not naturally coincide with the Landsat grid. We therefore calculated subsequent one-hectare plot NDVI averages as a weighted mean, where each pixel was weighted by the proportion of the plot area covered by the pixel.

#### 2.4. Ground-based plant phenology measurements

We carried out ground-based phenology monitoring in eight 2 m × 2 m plots (table S6), one adjacent to each one-hectare plot (mean distance = 23 m, max distance = 52 m). We placed the ground-based monitoring plots adjacent to the drone-based survey plots to minimise the effects of ecological disturbance and trampling in the drone survey plots caused by the repeat visits necessary for the ground-based monitoring. Within these plots, we monitored six individual plants from the most common species: *E. vaginatum*, *D. integrifolia*, *S. pulchra* and *A. latifolia* in tussock sedge tundra; *D. integrifolia*, *S. arctica* and *A. latifolia* in Dryas-vetch tundra. On each survey date, we measured the length of the longest leaf on each individual to the nearest millimetre. This approach is widely used in field-based phenology monitoring protocols [95], and will allow for NDVI to be directly related to phenological changes in plant traits. We conducted the ground-based surveys in tandem with the drone-based surveys where logistical possible, resulting in a dataset of 52 drone and ground survey pairs spread over 20 different dates. The majority of ground-based phenology surveys were carried out on the same day as the drone surveys (mean difference = 0.3 d, maximum difference = 3 d, table S7).

#### 2.5. Cross-sensor correspondence

To test cross-sensor correspondence, we first had to scale all datasets to a shared spatial grain and time-window. To achieve this, we first plotted the spatial mean NDVI for all one-hectare plots, time-points and available sensors (MODIS = single pixel, Landsat 8 = weighted mean) across both growing seasons (2016 and 2017). We then divided the two growing seasons into 22 consecutive seven-day blocks starting on the 1st of May each year. Next, we calculated the temporal mean of the spatial mean NDVI for each seven-day block for all plot and sensor combinations where data was available. We then identified all matching seven-day block and study plot combinations for each drone-satellite and satellite-satellite combination. We then tested cross-sensor correspondence by calculating Spearman's rank correlation and mean sensor-to-sensor difference in the plot means across the whole data set.

Additionally, we matched all drone and Sentinel-2 scenes, as well as all drone and Landsat 8 scenes that were less than two days apart. We resampled the red and near-infrared drone bands to the relevant Sentinel-2/Landsat 8 grids and calculated the NDVI. We restricted the analysis to Landsat 8 pixels fully contained within the study plots and reprojected the drone data from UTM 7 N to UTM 8 N using a bilinear reprojection where the Landsat 8 scenes were provided in this projection. Finally, we tested the predictive relationship between the resampled drone and satellite NDVI pixel-pairs for a random subsample of Sentinel pixels (10% of total,  $n = 700$ ) and all available Landsat 8 pixels ( $n = 198$ ) with Bayesian linear models (tables S8 and S9 for Sentinel-2, S10 and S11 for Landsat 8) using the MCMCglmm v.2.29 package [96].

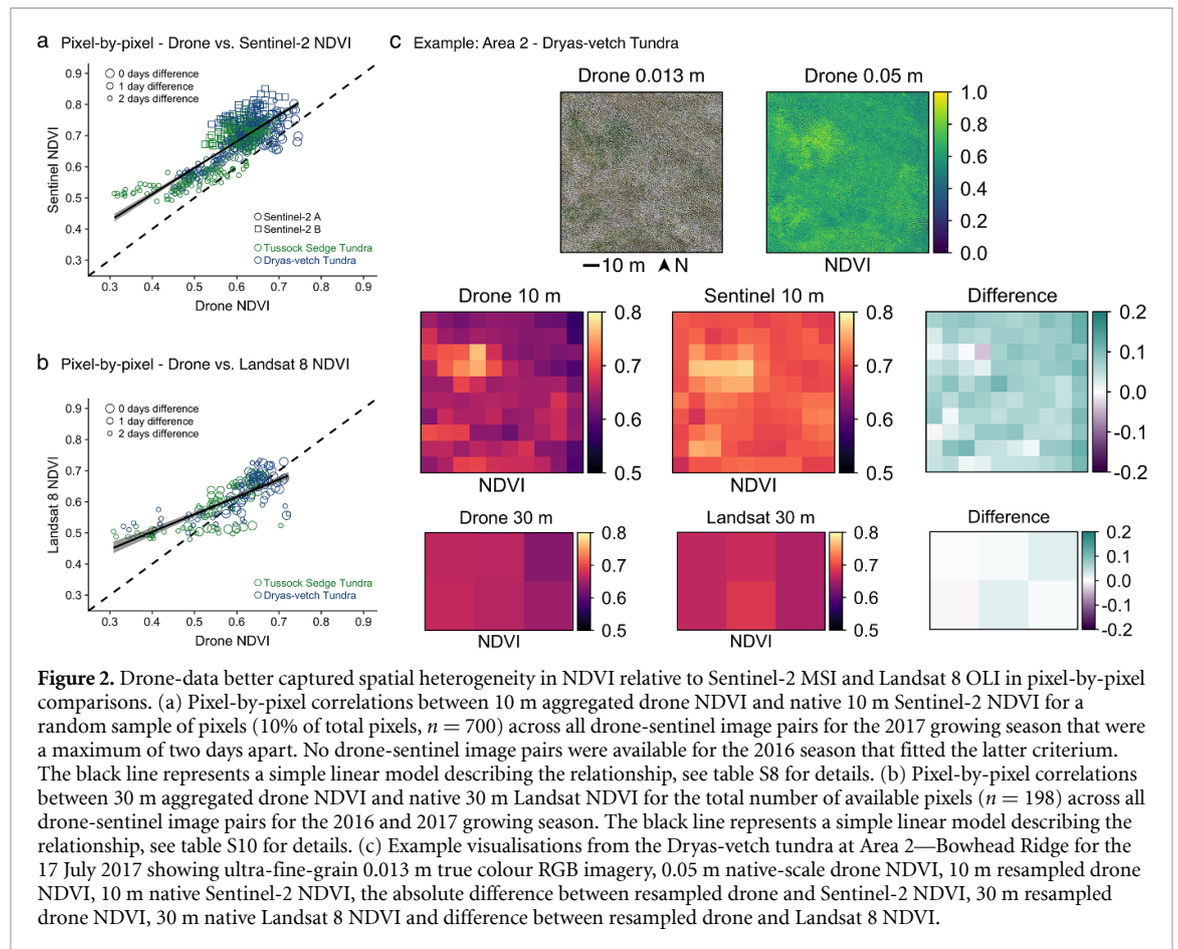
We used the 'resample' function of the R raster package v. 3.0–12 [97] for resampling from finer to coarser resolutions. The function first aggregates the smaller grid to the largest clean divisor of the larger grid using the mean and then, if required, resamples to the larger grid using bilinear interpolation. We also tested an alternative resampling approach by first resampling to a common resolution and grid of 0.5 m and then aggregating by mean, but found no qualitative differences in our results (figure S2). Further details about software and package versions used for raster manipulations and visualisations can be found in the supplementary materials.

#### 2.6. Spatial autocorrelation

To assess the spatial autocorrelation of variation in tundra greenness within the eight plots, we sampled variograms and fitted variogram models using the gstat v. 2.0–5 package [98, 99]. First, we pre-thinned the acquired drone-data by randomly sampling 5% of the ca. 4 million pixels of each orthomosaic. We then sampled variograms for all plots at the peak of the 2017 season (26 and 28 July) and fitted variogram models, letting the gstat algorithm select the best fit amongst spherical, exponential and Matern models. The only exception was Area 3 for which the closest available complete set of flights was on the 18 July 2017. To test conformity of the variograms across the season, we repeated the analysis for the surveys from the 26 June and 9 August 2017 for Area 1 and 2. No change in the variogram patterns were observed across the 2017 season and we therefore assume that our analysis is representative for the 2016 season also. All variograms were sampled with a bin width of 0.15 m from 0 to 15 m and a bin width of 3 m from 0 to 45 m.

#### 2.7. Grain size and phenology

We tested the influence of grain-size on observations of tundra greenness phenology by fitting simplified growing season curves to the raster stacks for each plot and season. We first resampled the drone bands



for all time-points to grids with grain sizes of 0.5, 1, 5, 10, 20 and 33.33 m. We then calculated the NDVI and fitted simple quadratic models to each pixel in the growing season stacks ( $y = ax^2 + bx + c$ , where  $x$  is the day of year and  $y$  the pixel NDVI,  $a$  the quadratic coefficient,  $b$  the linear coefficient and  $c$  the constant term). We found a strong negative correlation between the quadratic and linear coefficients of the models (figure S6), and therefore selected only the quadratic coefficient for further analysis. Additional details on model choice and analysis can be found in the method section of the supplementary materials.

## 2.8. Ground validation

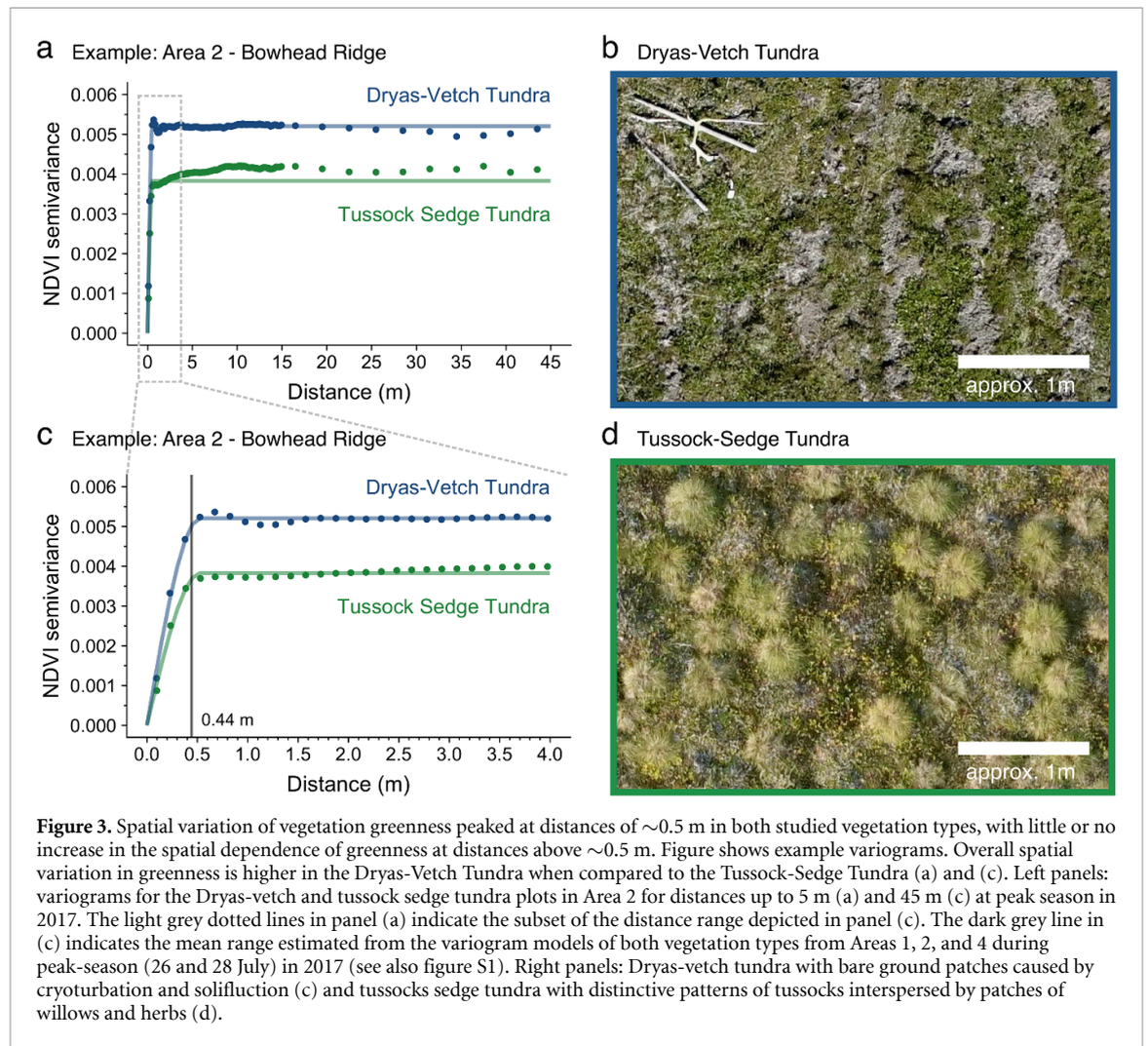
To test the correspondence between our ground-based phenology measurements and the drone observations, we derived time-series of the plot mean standardised longest leaf length (hereafter mean longest leaf length) for all species (using a z-score—centred data with a standard deviation of 1) and the drone-greenness for each  $2 \text{ m} \times 2 \text{ m}$  ground-based monitoring plot. See supplementary methods for details on how the leaf measurements were standardised. The drone-based plot mean NDVI values were then matched with the plot mean longest leaf length values from the closest ground-based survey date (table S7). We then calculated the Spearman's rank

correlation between mean NDVI and mean longest leaf length for each plot and season. We replicated the analysis using Sentinel-2 data where available (see supplementary materials). Finally, we also conducted a by-species version of the analysis using the by-species mean of the absolute longest leaf length for each  $2 \text{ m} \times 2 \text{ m}$  plot rather than the mean based on the standardised longest leaf lengths.

## 3. Results

### 3.1. Landscape greenness corresponded among sensors

Landscape greenness corresponded well among drone, Sentinel-2, Landsat 8 and MODIS across both the 2016 and 2017 growing seasons. Growing season curves of the mean NDVI for the one-hectare plots were similar (figure 1) and the plots' temporal (seven-day) mean NDVI values were highly correlated across sensors (Spearman's  $\rho > 0.59$ – $0.98$ , table S12). However, we observed a positive offset between the drone and satellite seven-day mean NDVI values for the plots. This offset ranged between 0.027 (Landsat 8) and 0.073 (Sentinel-2B) absolute NDVI and was consistently positive across satellites (table S13). The Landsat 8 offset of 0.027 fell within the range of the estimated error ( $\pm 0.03$ ) associated with the drone-derived mean NDVI for the study plots



determined in a previous study using the same survey method [81].

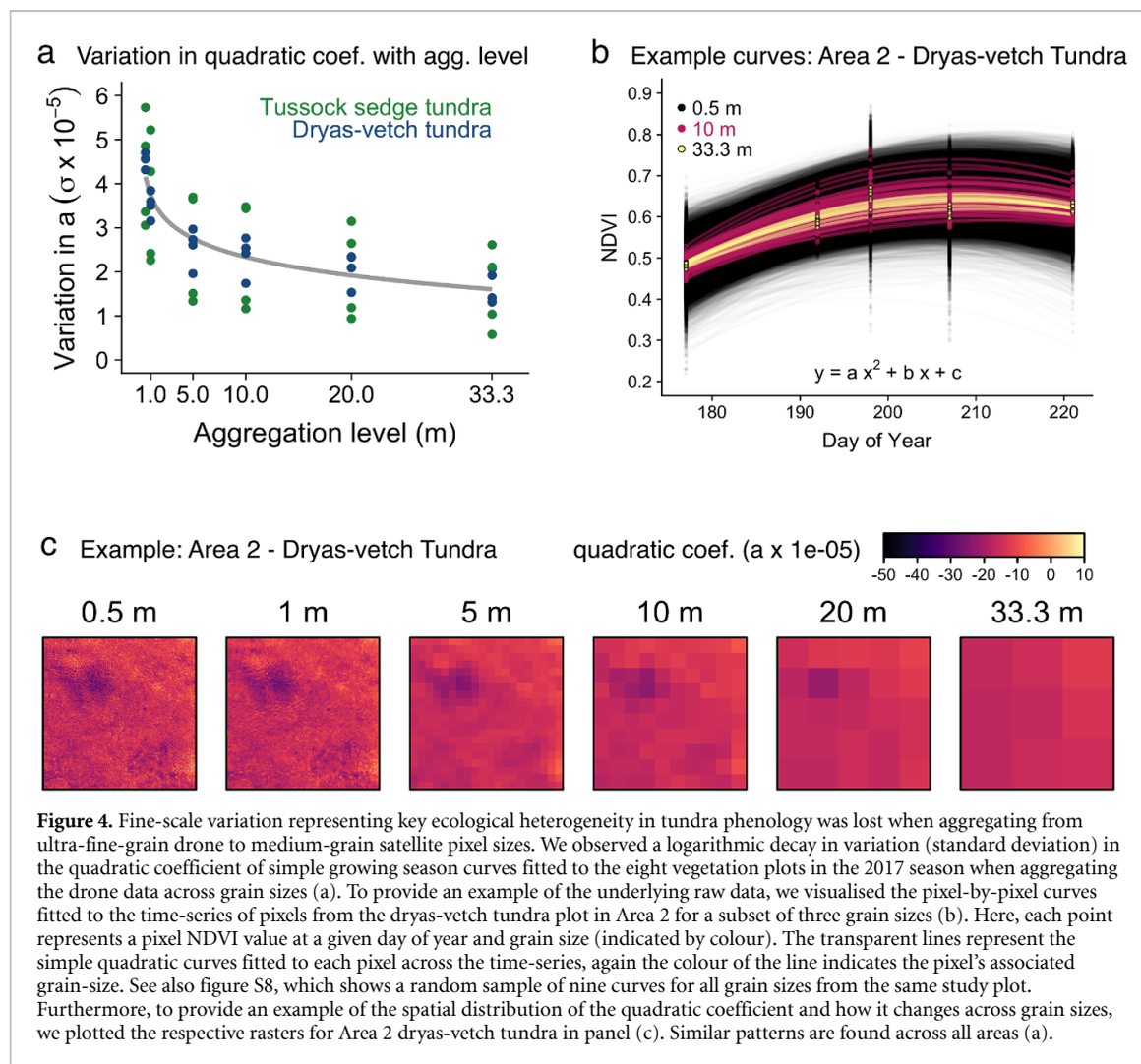
Resampled drone pixels (10 m and 30 m) and the corresponding spatially co-located Sentinel-2 and Landsat 8 pixels were highly correlated (marginal  $R^2 = 0.69$  and marginal  $R^2 = 0.58$  respectively, see figure 2 and tables S8 and S10). We found that vegetation type, the time-difference between satellite scene and drone data acquisition, and the specific Sentinel platform (Sentinel-2A/Sentinel-2B) influenced the relationship between Sentinel-2 pixel NDVI and drone-derived NDVI (marginal  $R^2 = 0.87$  see table S9). While the Sentinel platform (Sentinel-2A/Sentinel-2B) had the strongest impact on the intercept and the slope of the linear model, vegetation type and time-difference mainly influenced the slope, with time-difference having the smallest effect on slope and intercept overall (table S9). In contrast, we only detected a statistically meaningful effect for the time-difference between satellite and drone scene acquisition in the Landsat 8—drone pixel model (marginal  $R^2 = 0.70$ ); vegetation type did not have a statistically meaningful influence on this relationship (table S11).

### 3.2. Spatial variation in landscape greenness peaked at approx. 0.5 m

Spatial variability in the NDVI values associated with distance peaked at ranges below 0.5 meter (mean range 0.44 m) during the peak-season of 2017 (26–28 July). Little additional autocorrelation structure in the NDVI was found between pixel pairs for distances of up to 45 m (figure 3). This pattern was consistent across vegetation types in seven out of our eight plots (figures 3, S3 and S4). The only exception is the Dryas-vetch plot in Area 3, which showed the same patterns for distances below 10 m, but thereafter spatial variation steadily increased (figure S4). Peak variability (sill) in NDVI decreased as the growing season progressed (figure S5), and varied with vegetation type (figures 3, S3 and S4). Unexplained variability (nugget) was consistently low across all Areas (figures 3, S3 and S4).

### 3.3. Seasonal-variation was lost when aggregating to medium grain sizes

We observed a notable loss in the amount of seasonal variation in tundra greenness when aggregating



grain sizes from ultra-fine-grain drone to medium-grain satellite data. The loss was particularly pronounced at grain-sizes above 10 m—the grain size of Sentinel-2 MSI pixels (46.2%–63.9%) (figure 4). The variation in the quadratic coefficient of the simple growing season curves (figures 4(b) and S6) decayed logarithmically with grain size (figure 4(a)), while no change occurred in the mean tendency of the coefficient (figure S7). The quadratic and linear coefficients of the growing season curves were strongly correlated (Spearman's  $\rho = -0.999$ ), thus the same pattern holds true for the linear component of the curve fit (figure S6).

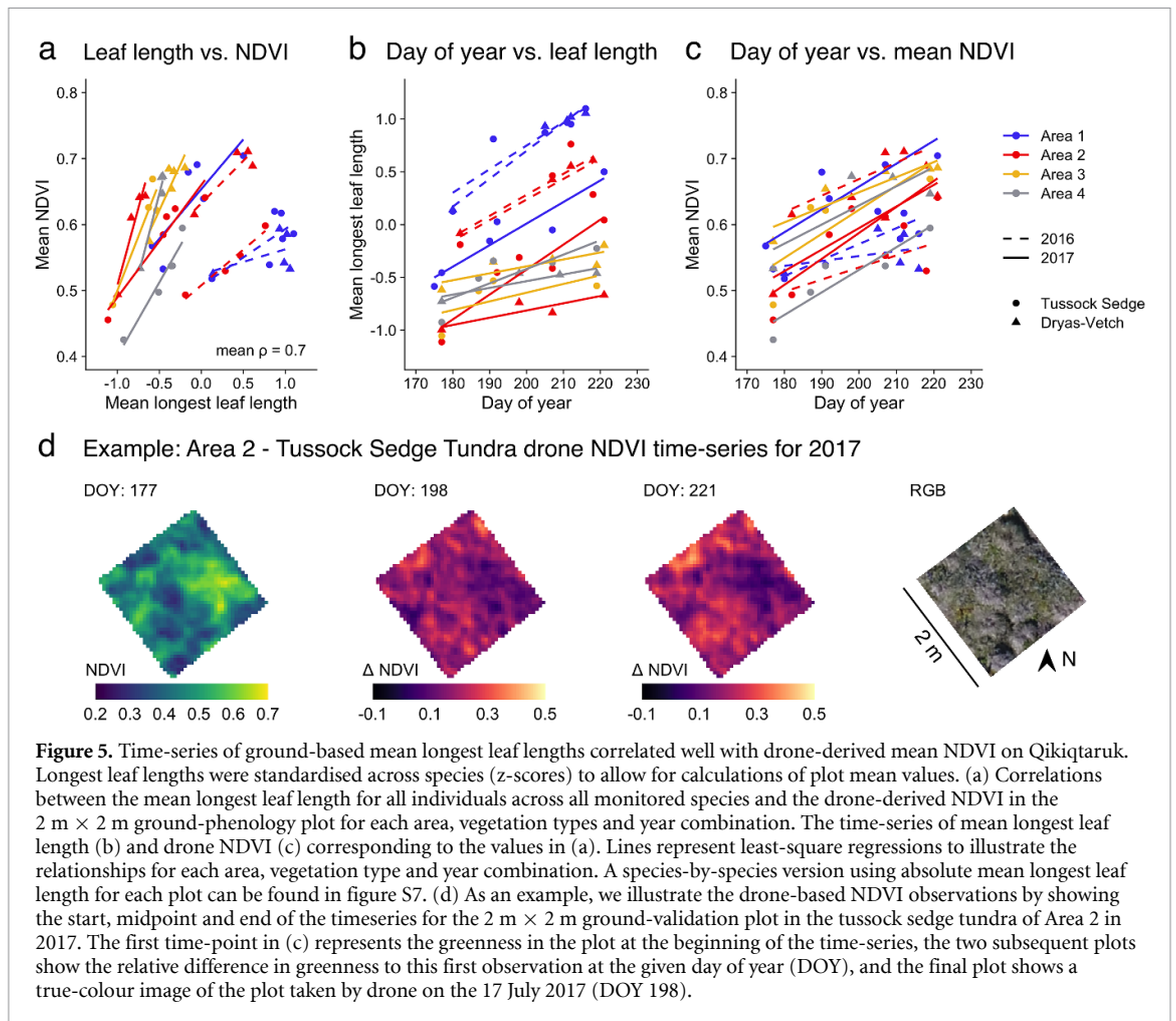
### 3.4. Drone-derived spectral greenness correlated well with leaf measurements

Drone-derived spectral greenness correlated well (mean  $\rho = 0.70$ ) with ground-based measurements of phenology for graminoids and deciduous plants across the growing season (figure 5). The Spearman's correlation coefficient of the plot mean longest leaf length and the mean drone-derived NDVI (mean  $\rho = 0.70$ , table S14 and figure 5) matched the by-species analysis based on absolute leaf lengths in

the ground-based phenology plots (mean  $\rho = 0.68$ , table S15 and figure S9). The graminoids and deciduous shrub species followed this mean tendency well across all time-series, while the partially-evergreen *D. integrifolia* showed mixed responses between plots and years (mean  $\rho = 0.22$ , figure S9). The drone-based time-series of greenness of the  $2\text{ m} \times 2\text{ m}$  ground-phenology plots highlight fine-scale differences in phenology such as the continuous greening of tussocks that was visible at the tussock sedge tundra plot in Area 2 (figure 5(c)). Sentinel-2 greenness of the ground-monitoring plots showed slightly weaker correlations (mean  $\rho = 0.58$ , figure S10) with the mean longest leaf length, but for this analysis no time-series of sufficient length were available for 2016 and peak-season observations in 2017 were limited.

## 4. Discussion

Our analysis of time-series of landscape greenness on Qikiqtaruk across scales highlights four main findings: 1) Measures of mean tendency in landscape greenness were consistent across sensors, but drone-derived NDVI values were lower than those



from Sentinel-2, Landsat 8 and MODIS products (figures 1 and 2). 2) The majority of variation in landscape greenness was contained at scales of around half-a-metre, and is thus not captured by medium-grain satellites such as Sentinel-2 (figure 3). 3) When aggregating growing season curves from ultra-fine-grain drone to medium-grain satellite pixel sizes, a notable amount (46.2%–63.9%) of variation in greenness phenology was lost (figure 4). 4) Drone-based measures of landscape greenness correlated well with ground-based measurements of leaf length (figure 5). Taken together, our results highlight that drone platforms and compact multispectral sensors can capture key ecological processes such as vegetation phenology and enable us to bridge the existing scale gap between satellite and ground-based monitoring in tundra ecosystems.

The correspondence between drone and satellite-derived NDVI has yet to be comprehensively tested across Arctic sites [13, 14]. Siewert and Olofson [14] also demonstrate cross-sensor agreement between drone- and satellite-derived NDVI from Arctic Sweden. While similar or higher levels of cross-sensor agreement have been observed in other natural and agricultural systems [14, 100, 101], some non-Arctic studies showed mixed or poor agreement [102–104].

Continued efforts in replicating these studies at different sites and systems are much needed to comprehensively evaluate cross-sensor correspondence in Arctic tundra systems and beyond.

We observed a small but consistent offset between drone- and satellite-derived NDVI that warrants further investigation. A similar offset has been detected in rice fields in Italy [103] and with spectroradiometer readings in ecologically similar tundra in Alaska [77], but see Siewert and Olofson [14] for a lack of offset in the more heterogeneous tundra of Arctic Sweden. Both technical and ecological factors could explain the offset. We were not able to conduct spectroradiometer readings coinciding with our drone surveys for on-the-ground comparisons. Technical reasons for the observed offset may include: atmospheric effects, differences in viewing geometries, sensor properties (e.g. band widths) and signal processing between drones and satellites (e.g. radiometric calibration), but also among different drone studies. Ecologically, the variation in land cover (especially the presence/absence of non-vegetative surfaces) or topography within a landscape may influence the correspondence between measures of vegetation greenness across scales due to non-linearities in how different patch sizes and cover types

are aggregated when measured with the NDVI [12, 105]. The high homogeneity of the survey plots on Qikiqtaruk likely contributes to the strong correlation between drone- and satellite-derived NDVI that we have observed. Yet, in our drone data fine-grain patterns of higher and lower NDVI within the landscape were evident, including bare-ground patches and areas of more productive vegetation in wetter parts of the landscape (figures 1–3). Non-linearities in the scaling of these patches could contribute to the offset between satellite and drone NDVI that we observed on Qikiqtaruk. Further research is needed to evaluate cross-sensor and cross-scale correspondence in NDVI and other vegetation indices across Arctic tundra systems.

We found that a plateau of spatial variation in tundra greenness occurred around 0.5 m, approximately the same width as biological and environmental patterning at this site. The *E. vaginatum* sedges that dominate the tussock sedge vegetation type typically have diameters of ~0.1–0.5 m (figure 3(b)) [106]. The tussock sedge vegetation type is underlain by ice-wedge polygons that when thawed create bands of wetter or drier plant communities with widths of ~0.5 m–3.0 m [107]. Dryas-vetch vegetation is often found on gentle sloping uplands where active layer disturbances such as cryoturbation and solifluction create characteristic bare-ground patches perpendicular to the slope [85] with dimensions of ~0.3 m–0.5 m width and ~0.3–1.0 m length (figure 3(b)). We expect that spatial variation would increase with distances beyond the one-hectare extents of our plots as more topographic diverse terrain is encountered and vegetation type transitions are reached. Topography is a key proxy for many processes that structure heterogeneity in tundra vegetation [108–110] and the plots were selected for little topographic variation to allow us to isolate specific effects of land cover on scaling of greenness patterns from topography. The plot with the highest elevational range (Area 3—Dryas-vetch tundra: 8.7 m) showed a small but steady increase in spatial variation in distance classes above 10 m (figure S4). Our findings illustrate that on Qikiqtaruk, grain sizes of 0.5 m or less are required to capture key spatial variation in vegetation greenness.

In our study, ecological information was lost when upscaling from ultra-fine-grain (~0.05 m) drone to moderate grain (~10–30 m) satellite resolutions. Even the most recent generation of freely-available multispectral satellite products can be limited in their ability to capture fine-grain ecological processes of tundra vegetation change [13]. Information transfer during upscaling leads to the loss of more information in tundra ecosystems compared to other biomes [14, 111] as land cover and vegetation structure are fragmented at finer scales [112]. However, exactly how spatial aggregation influences the loss in observed ecological variability across the diversity of Arctic landscapes remains

poorly quantified [11]. Yet, this variability is critical to understanding climate-driven changes in vegetation phenology [35, 36, 88], plant-pollinator interactions [113], and trophic interactions [114]. With fine-grain observations, we were able to detect a subtle decrease in the magnitude of the spatial variability in landscape-level phenology as the growing season progressed (figure S5), while aggregation to moderate satellite grains obscured both the magnitude and timing of phenological heterogeneity (figure 4). Thus, time-series of fine-grain remotely-sensed observations will be critical for answering key research questions about tundra ecosystem functioning in a warming Arctic [115].

Our results indicate that drone-based greenness time-series captured variation in leaf-growth of deciduous tundra plant species at the plot level. We demonstrate how drones can be used to measure variation in tundra plant phenology of metre-scale patches at landscape extents. Drones have been successfully used to monitor phenology of individual plants (trees) in temperate forest ecosystems [116–118], and our ability to detect sub-decimeter variability in our study indicates that individual plant-level phenology monitoring with drones could also be carried out in the tundra. Future studies that quantify plant growth or phenology events such as leaf emergence and flowering across the landscape could provide key information on resource availability for plant-consumer interactions [113, 114]. Our findings also highlight known limitations of NDVI to track phenology in evergreens or other non-deciduous taxa (*D. integrifolia*, figure S9), suggesting that tests of alternative vegetation index—plant growth relationships [118] are needed to capture variation in plant metabolic activity of tundra evergreen and moss species within the growing season. Combining drone-based time-series with observations from phenocams, satellite and ground-based study plots has the potential to revolutionise our understanding of landscape-scale phenology [13] by moving beyond the previously small samples of individuals monitored in the Arctic tundra [36, 37, 39, 119].

The collection of multispectral drone time-series in Arctic ecosystems has limitations and challenges. Recent studies have discussed challenges with radiometric consistency and repeatability when using compact multispectral drone sensors [81, 120, 121]. Due to logistical constraints, we were not able to always conduct surveys under optimal conditions due to sun angle or cloud cover, nor as frequently as planned due to wind or precipitation (table S2), which likely introduced bias and/or noise into our drone data (e.g. figure 4(b)). Site access limitations meant that we could not capture spring and autumn on Qikiqtaruk. As an early-generation multispectral drone sensor, the Parrot Sequoia was tailored for deriving the NDVI, which despite being the legacy

workhorse of tundra remote-sensing has limitations [11, 13]. In particular, NDVI can be confounded by moisture and surface water [11, 73, 122], complicating interpretation in wet tundra, particularly at fine-grain sizes. However, the rapid technological development of drones and sensors, as well as further consolidation and standardisation of methods [123], will allow for pan-Arctic syntheses of fine-grain data to resolve the uncertainty and complexity of Arctic greening patterns trends [13, 14, 81] (see also the High Latitude Drone Ecology Network—(<https://arcticdrones.org/>)).

Our study demonstrates that drones can fill the scale-gap between satellite and field studies of terrestrial Arctic vegetation change. Rather than investigating and explaining patterns at scales pre-defined by satellite datasets or field-based networks, researchers can use drones to identify scale-domains that are most closely associated with the ecological processes of interest. Field ecologists can now combine scaling theory provided by the remote sensing community [74, 124–127] with observations at scales and temporal intervals that allow for the testing of hypotheses about the mechanisms that drive landscape-level ecological change. Drone imagery will also allow the remote sensing community to track the effects of sub-pixel heterogeneity on satellite products down to the grain of individual plants and communities [14], which have been long studied by field-based monitoring networks, like the International Tundra Experiment [75]. Only by improving our understanding of how ecologically important information is captured across grain sizes can we reduce uncertainties in the medium- and coarse-grain satellite observation that feed into Earth system models and shape their predictions [4, 8]. Fine-scale remote sensing from drones and aircraft therefore provide key tools for disentangling the drivers behind the greening of the Arctic [14, 79, 115].

## 5. Conclusions

Novel remote-sensing technologies such as drones now allow us to study ecological variation in landscapes continuously across scales. Fine-grain ecological observations are of particular importance where variation in plant growth happens at very small spatial scales such as in tundra ecosystems [13, 71]. The peak in spatial variation we found at distances of  $\sim 0.5$  m in the plots on Qikiqtaruk demonstrates the grain size at which phenological information within the plant communities is best captured at this site. We show that key ecological information is lost when observing the tundra at even decimeter or coarser scales, such as those of medium grain satellites ( $\sim 10$ – $30$  m). Despite the methodological challenges of collecting multispectral drone imagery in remote environments [81], our time-series of vegetation greenness correlated well with ground-based measurements of

leaf growth in the validation plots. Drones now enable studies that fill the scale gaps between satellite and ground-based observations, and therefore improve our ability to identify the key drivers of vegetation change and project climate change impacts and feedbacks in the tundra biome.

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### Author contributions

JJA and IMS conceived the study with input from JTK and AMC. JJA carried out data processing and analysis. JJA and IMS led the drone and ground-validation field work in 2016. AMC led the drone field surveys in 2017 with input from JTK. GD led the ground-validation for 2017 with input from JTK. JJA, IMS and JTK wrote the manuscript with input from AMC and GD. IMS supervised and acquired funding for the research.

### Data availability statement

The data and code that support the findings of this study are openly available at the following URL/DOI: ([https://github.com/jakobjassmann/qhi\\_phen\\_ts](https://github.com/jakobjassmann/qhi_phen_ts)).

### ORCID iDs

Jakob J Assmann  <https://orcid.org/0000-0002-3492-8419>

Isla H Myers-Smith  <https://orcid.org/0000-0002-8417-6112>

Jeffrey T Kerby  <https://orcid.org/0000-0002-2739-9096>

Andrew M Cunliffe  <https://orcid.org/0000-0002-8346-4278>

Gergana N Daskalova  <https://orcid.org/0000-0002-5674-5322>

### References

- [1] Levin S A 1992 The problem of pattern and scale in ecology: the Robert H. MacArthur award lecture *Ecology* **73** 1943–67
- [2] Anderson C B 2018 Biodiversity monitoring, earth observations and the ecology of scale *Ecol. Lett.* **21** 1572–85
- [3] Estes L *et al* 2018 The spatial and temporal domains of modern ecology *Nat. Ecol. Evol.* **2** 819
- [4] Chapin F S *et al* 2005 Role of land-surface changes in arctic summer warming *Science* **310** 657–60
- [5] Loranty M M and Goetz S J 2012 Shrub expansion and climate feedbacks in Arctic tundra *Environ. Res. Lett.* **7** 011005
- [6] Pearson R G, Phillips S J, Loranty M M, Beck P S A, Damoulas T, Knight S J and Goetz S J 2013 Shifts in Arctic vegetation and associated feedbacks under climate change *Nat Clim Change*. **3** 673–7
- [7] Richardson A D, Keenan T F, Migliavacca M, Ryu Y, Sonnentag O and Toomey M 2013 Climate change, phenology, and phenological control of vegetation feedbacks to the climate system *Agric. Forest Meteorol.* **169** 156–73
- [8] Ernakovich J G, Hopping K A, Berdanier A B, Simpson R T, Kachergis E J, Steltzer H and Wallenstein M D 2014 Predicted responses of arctic and alpine ecosystems to altered seasonality under climate change *Glob. Change Biol.* **20** 3256–3269
- [9] Diepstraten R A E, Jessen T D, Fauvelle C M D and Musiani M M 2018 Does climate change and plant phenology research neglect the Arctic tundra? *Ecosphere* **9** e02362
- [10] Metcalfe D B *et al* 2018 Patchy field sampling biases understanding of climate change impacts across the Arctic *Nat. Ecol. Evol.* **2** 1443
- [11] Beamish A *et al* 2020 Recent trends and remaining challenges for optical remote sensing of Arctic tundra vegetation: a review and outlook *Remote Sens. Environ.* **246** 111872
- [12] Stoy P C, Williams M, Disney M, Prieto-Blanco A, Huntley B, Baxter R and Lewis P 2009 Upscaling as ecological information transfer: a simple framework with application to Arctic ecosystem carbon exchange *Landsc Ecol.* **24** 971–86
- [13] Myers-Smith I H *et al* 2020 Complexity revealed in the greening of the Arctic *Nat Clim Change*. **10** 106–17
- [14] Siewert M B and Olofsson J 2020 Scale-dependency of Arctic ecosystem properties revealed by UAV *Environ. Res. Lett.* **15** 094030
- [15] Myneni R B, Keeling C D, Tucker C J, Asrar G and Nemani R R 1997 Increased plant growth in the northern high latitudes from 1981 to 1991 *Nature* **386** 698–702
- [16] Tucker C J, Slayback D A, Pinzon J E, Los S O, Myneni R B and Taylor M G 2001 Higher northern latitude normalized difference vegetation index and growing season trends from 1982 to 1999 *Int. J. Biometeorol.* **45** 184–90
- [17] Bhatt U S *et al* 2010 Circumpolar arctic tundra vegetation change is linked to sea ice decline *Earth Interact.* **14** 1–20
- [18] Guay K C, Beck P S A, Berner L T, Goetz S J, Baccini A and Buermann W 2014 Vegetation productivity patterns at high northern latitudes: a multi-sensor satellite data assessment *Glob. Change Biol.* **20** 3147–58
- [19] Zhu Z *et al* 2016 Greening of the Earth and its drivers *Nat Clim Change*. **6** 791–5
- [20] Keenan T F and Riley W J 2018 Greening of the land surface in the world's cold regions consistent with recent warming *Nat Clim Change*. **8** 825–8
- [21] Zeng H, Jia G and Epstein H 2011 Recent changes in phenology over the northern high latitudes detected from multi-satellite data *Environ. Res. Lett.* **6** 045508
- [22] Zeng H, Jia G and Forbes B C 2013 Shifts in Arctic phenology in response to climate and anthropogenic factors as detected from multiple satellite time series *Environ. Res. Lett.* **8** 035036
- [23] Zhao J, Zhang H, Zhang Z, Guo X, Li X and Chen C 2015 Spatial and temporal changes in vegetation phenology at middle and high latitudes of the northern hemisphere over the past three decades *Remote Sens.* **7** 10973–95
- [24] Potter C and Alexander O 2020 Changes in vegetation phenology and productivity in Alaska over the past two decades *Remote Sens.* **12** 1546
- [25] Tucker C J 1979 Red and photographic infrared linear combinations for monitoring vegetation *Remote Sens. Environ.* **8** 127–50
- [26] Myneni R B, Tucker C J, Asrar G and Keeling C D 1998 Interannual variations in satellite-sensed vegetation index data from 1981 to 1991 *J. Geophys. Res. Atmos.* **103** 6145–60
- [27] Elmendorf S C *et al* 2012 Plot-scale evidence of tundra vegetation change and links to recent summer warming *Nat Clim Change*. **2** 453–7
- [28] Tape K D, Sturm M and Racine C H 2006 The evidence for shrub expansion in northern Alaska and the Pan-Arctic *Glob. Change Biol.* **12** 686–702
- [29] Myers-Smith I H *et al* 2011 Shrub expansion in tundra ecosystems: dynamics, impacts and research priorities *Environ. Res. Lett.* **6** 045509
- [30] Tape K, Hallinger M, Welker J and Ruess R 2012 Landscape heterogeneity of shrub expansion in Arctic Alaska *Ecosystems* **15** 711–24
- [31] García Criado M, Myers-Smith I H, Bjorkman A D, Lehmann C E R and Stevens N 2020 Woody plant

- encroachment intensifies under climate change across tundra and savanna biomes *Glob. Ecol. Biogeogr.* **29** 925–43
- [32] Bjorkman A D *et al* 2018 Plant functional trait change across a warming tundra biome *Nature* **562** 57–62
- [33] Høye T T, Post E, Meltofte H, Schmidt N M and Forchhammer M C 2007 Rapid advancement of spring in the high arctic *Curr. Biol.* **17** R449–51
- [34] Kerby J T and Post E 2013 Advancing plant phenology and reduced herbivore production in a terrestrial system associated with sea ice decline *Nat. Commun.* **4** 2514
- [35] Post E, Kerby J, Pedersen C and Steltzer H 2016 Highly individualistic rates of plant phenological advance associated with arctic sea ice dynamics *Biol. Lett.* **12** 20160332
- [36] Assmann J J *et al* 2019 Local snow melt and temperature—but not regional sea ice—explain variation in spring phenology in coastal Arctic tundra *Glob. Change Biol.* **25** 2258–74
- [37] Oberbauer S F *et al* 2013 Phenological response of tundra plants to background climate variation tested using the international tundra experiment *Philos. Trans. R. Soc. B* **368** 20120481 (<http://rstb.royalsocietypublishing.org/content/368/1624/20120481>)
- [38] Bjorkman A D, Elmendorf S C, Beamish A L, Vellend M and Henry G H R 2015 Contrasting effects of warming and increased snowfall on Arctic tundra plant phenology over the past two decades *Glob. Change Biol.* **21** 4651–61
- [39] Prev y J S *et al* 2019 Warming shortens flowering seasons of tundra plant communities *Nat. Ecol. Evol.* **3** 45
- [40] Jia G J, Epstein H E and Walker D A 2006 Spatial heterogeneity of tundra vegetation response to recent temperature changes *Glob. Change Biol.* **12** 42–55
- [41] Reynolds M K, Comiso J C, Walker D A and Verbyla D 2008 Relationship between satellite-derived land surface temperatures, arctic vegetation types, and NDVI *Remote Sens. Environ.* **112** 1884–94
- [42] Jia G J, Epstein H E and Walker D A 2009 Vegetation greening in the Canadian arctic related to decadal warming *J. Environ. Monit.* **11** 2231
- [43] Bhatt U S *et al* 2013 Recent declines in warming and vegetation greening trends over Pan-Arctic tundra *Remote Sens.* **5** 4229–54
- [44] Xu L *et al* 2013 Temperature and vegetation seasonality diminishment over northern lands *Nat. Clim. Change* **3** 581–586
- [45] Piao S *et al* 2014 Evidence for a weakening relationship between interannual temperature variability and northern vegetation activity *Nat. Commun.* **5** 5018
- [46] Vickers H *et al* 2016 Changes in greening in the high Arctic: insights from a 30 year AVHRR max NDVI dataset for Svalbard *Environ. Res. Lett.* **11** 105004
- [47] Walker D A *et al* 2009 Spatial and temporal patterns of greenness on the Yamal Peninsula, Russia: interactions of ecological and social factors affecting the Arctic normalized difference vegetation index *Environ. Res. Lett.* **4** 045004
- [48] Macias-Fauria M, Forbes B C, Zetterberg P and Kumpula T 2012 Eurasian Arctic greening reveals teleconnections and the potential for structurally novel ecosystems *Nat. Clim. Change.* **2** 613–8
- [49] Post E 2013 Erosion of community diversity and stability by herbivore removal under warming *Proc. R. Soc. B* **280** 20122722
- [50] Bhatt U S *et al* 2014 Implications of Arctic sea ice decline for the Earth system *Annu. Rev. Environ. Resour.* **39** 57–89
- [51] Fauchald P, Park T, Tømmervik H, Myneni R and Hausner V H 2017 Arctic greening from warming promotes declines in caribou populations *Sci. Adv.* **3** e1601365
- [52] Elmendorf S C *et al* 2015 Experiment, monitoring, and gradient methods used to infer climate change effects on plant communities yield consistent patterns *Proc. Natl Acad. Sci.* **112** 448–52
- [53] Pattison R R, Jorgenson J C, Reynolds M K and Welker J M 2015 Trends in NDVI and tundra community composition in the Arctic of NE Alaska between 1984 and 2009 *Ecosystems* **18** 707–19
- [54] Piao S *et al* 2015 Leaf onset in the northern hemisphere triggered by daytime temperature *Nat. Commun.* **6** 6911
- [55] Liu Q *et al* 2016 Delayed autumn phenology in the Northern Hemisphere is related to change in both climate and spring phenology *Glob. Change Biol.* **22** 3702–11
- [56] Semenchuk P R, Gillespie M A K, Rumpf S B, Baggesen N, Elberling B and Cooper E J 2016 High Arctic plant phenology is determined by snowmelt patterns but duration of phenological periods is fixed: an example of periodicity *Environ. Res. Lett.* **11** 125006
- [57] Prev y J *et al* 2017 Greater temperature sensitivity of plant phenology at colder sites: implications for convergence across northern latitudes *Glob. Change Biol.* **3** 2660–71
- [58] Jia G J, Epstein H E and Walker D A 2003 Greening of arctic Alaska, 1981–2001 *Geophys. Res. Lett.* **30** HLS 3–1
- [59] Fraser R H, Olthof I, Carri re M, Deschamps A and Pouliot D 2011 Detecting long-term changes to vegetation in northern Canada using the Landsat satellite image archive *Environ. Res. Lett.* **6** 045502
- [60] Ju J and Masek J G 2016 The vegetation greenness trend in Canada and US Alaska from 1984–2012 Landsat data *Remote Sens. Environ.* **176** 1–16
- [61] Reynolds M K, Walker D A, Verbyla D and Munger C A 2013 Patterns of change within a Tundra Landscape: 22-year Landsat NDVI trends in an area of the northern foothills of the Brooks range, Alaska *Arct. Antarct. Alp. Res.* **45** 249–60
- [62] Miles V V and Esau I 2016 Spatial heterogeneity of greening and browning between and within bioclimatic zones in northern West Siberia *Environ. Res. Lett.* **11** 115002
- [63] Lara M J, Nitze I, Grosse G, Martin P and McGuire A D 2018 Reduced arctic tundra productivity linked with landform and climate change interactions *Sci. Rep.* **8** 2345
- [64] Thompson J A and Koenig L S 2018 Vegetation phenology in Greenland and links to cryospheric change *Ann. Glaciol.* **59** 59–68
- [65] Phoenix G K and Bjerke J W 2016 Arctic browning: extreme events and trends reversing arctic greening *Glob. Change Biol.* **22** 2960–2
- [66] National Academies of Sciences E 2019 Understanding Northern latitude vegetation greening and browning: proceedings of a workshop (available at: (<https://www.nap.edu/catalog/25423/understanding-northern-latitude-vegetation-greening-and-browning-proceedings-of-a>) (Accessed 21 October 2019)
- [67] Frost G V *et al* 2019 Arctic report card: update for 2019 - Tundra greenness Arctic Program (available at: (<http://www.arctic.noaa.gov/Report-Card/Report-Card-2015/ArtMID/5037/ArticleID/221/Tundra-Greenness>) (Accessed 23 February 2019)
- [68] Blok D, Schaepman-Strub G, Bartholomeus H, Heijmans M M P D, Maximov T C and Berendse F 2011 The response of Arctic vegetation to the summer climate: relation between shrub cover, NDVI, surface albedo and temperature *Environ. Res. Lett.* **6** 035502
- [69] Reynolds M K, Walker D A, Epstein H E, Pinzon J E and Tucker C J 2012 A new estimate of tundra-biome phytomass from trans-Arctic field data and AVHRR NDVI *Remote Sens. Lett.* **3** 403–11
- [70] Berner L T, Jantz P, Tape K D and Goetz S J 2018 Tundra plant above-ground biomass and shrub dominance mapped across the North slope of Alaska *Environ. Res. Lett.* **13** 035002
- [71] Cunliffe A M, Assmann J J, Daskalova G, Kerby J T and Myers-Smith I H 2020 Aboveground biomass corresponds strongly with drone-derived canopy height but weakly with greenness (NDVI) in a shrub tundra landscape *Environ. Res. Lett.* (<https://doi.org/10.1088/1748-9326/aba470>)
- [72] Stow D 2004 Remote sensing of vegetation and land-cover change in arctic tundra ecosystems *Remote Sens. Environ.* **89** 281–308

- [73] Bartsch A *et al* 2020 Feasibility of tundra vegetation height retrieval from Sentinel-1 and Sentinel-2 data *Remote Sens. Environ.* **237** 111515
- [74] Woodcock C E and Strahler A H 1987 The factor of scale in remote sensing *Remote Sens. Environ.* **21** 311–32
- [75] Henry G H R and Molau U 1997 Tundra plants and climate change: the international tundra experiment (ITEX) *Glob. Change Biol.* **3** 1–9
- [76] Beck P S A, Jönsson P, Høgda K-A, Karlsen S R, Eklundh L and Skidmore A K 2007 A ground-validated NDVI dataset for monitoring vegetation dynamics and mapping phenology in Fennoscandia and the Kola peninsula *Int. J. Remote Sens.* **28** 4311–30
- [77] Gamon J A, Huemmrich K F, Stone R S and Tweedie C E 2013 Spatial and temporal variation in primary productivity (NDVI) of coastal Alaskan tundra: decreased vegetation growth following earlier snowmelt *Remote Sens. Environ.* **129** 144–53
- [78] Kerby J T 2015 Phenology in a changing Arctic: linking trophic interactions across scales Sep 14 (available at: (<https://etda.libraries.psu.edu/catalog/26992>)) (Accessed 19 October 2018)
- [79] Anderson K and Gaston K J 2013 Lightweight unmanned aerial vehicles will revolutionize spatial ecology *Front Ecol. Environ.* **11** 138–46
- [80] Klosterman S and Richardson A D 2017 Observing spring and fall phenology in a deciduous forest with aerial drone imagery *Sensors.* **17** 2852
- [81] Assmann J J, Kerby J T, Cunliffe A M and Myers-Smith I H 2018 Vegetation monitoring using multispectral sensors — best practices and lessons learned from high latitudes *J. Unmanned Veh. Syst.* **7** 54–75
- [82] Klosterman S, Hufkens K and Richardson A D 2018 Later springs green-up faster: the relation between onset and completion of green-up in deciduous forests of North America *Int. J. Biometeorol.* **62** 1645–55
- [83] Myers-Smith I H and Hik D S 2013 Shrub canopies influence soil temperatures but not nutrient dynamics: an experimental test of tundra snow–shrub interactions *Ecol. Evol.* **3** 3683–700
- [84] Smith C, Kennedy C, Hargrave A and McKenna K 1989 Soil and vegetation of Herschel Island. Whitehorse, Yukon Territory, Canada: agriculture Canada Report No.: No 1
- [85] Obu J, Lantuit H, Myers-Smith I, Heim B, Wolter J and Fritz M 2015 Effect of terrain characteristics on soil organic carbon and total nitrogen stocks in soils of Herschel Island, Western Canadian Arctic *Permafrost. Periglacial Process.* **28** 92–107
- [86] Myers-Smith I H *et al* 2011 Expansion of canopy-forming willows over the twentieth century on Herschel Island, Yukon Territory, Canada *AMBIO* **40** 610–23
- [87] Raynolds M K *et al* 2019 A raster version of the circumpolar arctic vegetation map (CAVM) *Remote Sens. Environ.* **232** 111297
- [88] Myers-Smith I H *et al* 2019 Eighteen years of ecological monitoring reveals multiple lines of evidence for tundra vegetation change *Ecol. Monogr.* **89** e01351
- [89] Becker R A, Wilks A R, Brownrigg R, Minka T P and Deckmyn A 2018 maps: draw geographical maps (available at: (<https://CRAN.R-project.org/package=maps>))
- [90] Becker R A, Wilks A R and Brownrigg R 2018 mapdata: extra map databases (available at: (<https://CRAN.R-project.org/package=mapdata>))
- [91] Assmann J J, Myers-Smith I H, Kerby J T, Cunliffe A M and Daskalova G N 2020 Qikiqtaruk Phenology Time-Series code repository ([https://github.com/jakobjassmann/qhi\\_phen\\_ts](https://github.com/jakobjassmann/qhi_phen_ts))
- [92] Didan K 2015 MOD13Q1 MODIS/Terra Vegetation Indices 16-Day L3 Global 250m SIN Grid V006 NASA EOSDIS Land Processes DAAC (available at: (<https://doi.org/10.5067/MODIS/MOD13Q1.006>)) (Accessed 22 October 2018)
- [93] Gorelick N, Hancher M, Dixon M, Ilyushchenko S, Thau D and Moore R 2017 Google earth engine: planetary-scale geospatial analysis for everyone *Remote Sens. Environ.* **202** 18–27
- [94] Mueller-Wilm U 2017 Sen2Cor software release note: ref.: S2-PDGS-MPC-L2A-SRN-V2.4.0 (available at: ([http://step.esa.int/thirdparties/sen2cor/2.4.0/Sen2Cor\\_240\\_Documentation\\_PDF/S2-PDGS-MPC-L2A-SRN-V2.4.0.pdf](http://step.esa.int/thirdparties/sen2cor/2.4.0/Sen2Cor_240_Documentation_PDF/S2-PDGS-MPC-L2A-SRN-V2.4.0.pdf))) (Accessed 22 Oct 2018)
- [95] Molau U and Mølgaard P 1996 International tundra experiment manual (available at: (<https://www.gvsu.edu/itex/library-8.htm>))
- [96] Hadfield J D 2010 MCMC methods for multi-response generalized linear mixed models: the **MCMCglmm** R package *J. Stat. Softw.* **33** 1–22
- [97] Hijmans R J 2020 raster: geographic data analysis and modeling (available at: (<https://CRAN.R-project.org/package=raster>))
- [98] Pebesma E J 2004 Multivariable geostatistics in S: the gstat package *Comput. Geosci.* **30** 683–91
- [99] Gräler B, Pebesma E and Heuvelink G 2016 Spatio-Temporal Interpolation using gstat *R J.* **8** 204–18
- [100] Fernández-Guisuraga J M, Sanz-Abianedo E, Suárez-Seoane S and Calvo L 2018 Using unmanned aerial vehicles in postfire vegetation survey campaigns through large and heterogeneous areas: opportunities and challenges *Sensors.* **18** 586
- [101] Fawcett D *et al* 2020 Multi-scale evaluation of drone-based multispectral surface reflectance and vegetation indices in operational conditions *Remote Sens.* **12** 514
- [102] Matese A, Toscano P, Di Gennaro S F, Genesio L, Vaccari F P, Primicerio J, Belli C, Zaldei A, Bianconi R and Gioli B 2015 Intercomparison of UAV, aircraft and satellite remote sensing platforms for precision viticulture *Remote Sens.* **7** 2971–90
- [103] Franzini M, Ronchetti G, Sona G and Casella V 2019 Geometric and radiometric consistency of parrot sequoia multispectral imagery for precision agriculture applications *Appl. Sci.* **9** 5314
- [104] Khaliq A, Comba L, Biglia A, Ricauda Aimonino D, Chiaberge M and Gay P 2019 Comparison of satellite and UAV-based multispectral imagery for vineyard variability assessment *Remote Sens.* **11** 436
- [105] Riihimäki H, Luoto M and Heiskanen J 2019 Estimating fractional cover of tundra vegetation at multiple scales using unmanned aerial systems and optical satellite data *Remote Sens. Environ.* **224** 119–32
- [106] Mark A F, Fetcher N, Shaver G R and Chapin III F S 1985 Estimated ages of mature tussocks of eriophorum vaginatum along a latitudinal gradient in central Alaska U.S.A. *Arct. Alp. Res.* **17** 1–5
- [107] Fritz M *et al* 2016 Holocene ice-wedge polygon development in northern Yukon permafrost peatlands (Canada) *Quat. Sci. Rev.* **147** 279–97
- [108] Billings W D and Bliss L C 1959 An alpine snowbank environment and its effects on vegetation, plant development, and productivity *Ecology* **40** 388–97
- [109] Suvanto S, Roux P C L and Luoto M 2014 Arctic-alpine vegetation biomass is driven by fine-scale abiotic heterogeneity *Geogr. Ann. Ser. Phys. Geogr.* **96** 549–60
- [110] Riihimäki H, Heiskanen J and Luoto M 2017 The effect of topography on arctic-alpine aboveground biomass and NDVI patterns *Int. J. Appl. Earth Obs. Geoinf.* **56** 44–53
- [111] Stoy P C, Williams M, Spadavecchia L, Bell R A, Prieto-Blanco A, Evans J G and van Wijk M T 2009 Using information theory to determine optimum pixel size and shape for ecological studies: aggregating land surface characteristics in arctic ecosystems *Ecosystems* **12** 574–89
- [112] Virtanen T and Ek M 2014 The fragmented nature of tundra landscape *Int. J. Appl. Earth Obs. Geoinf.* **27** 4–12

- [113] Høye T T, Post E, Schmidt N M, Trøjelsgaard K and Forchhammer M C 2013 Shorter flowering seasons and declining abundance of flower visitors in a warmer Arctic *Nat Clim Change*. **3** 759–63
- [114] Armstrong J B, Takimoto G, Schindler D E, Hayes M M and Kauffman M J 2016 Resource waves: phenological diversity enhances foraging opportunities for mobile consumers *Ecology* **97** 1099–112
- [115] Miller C E et al 2019 An overview of ABoVE airborne campaign data acquisitions and science opportunities *Environ. Res Lett.* **14** 080201
- [116] Klosterman S et al 2018 Fine-scale perspectives on landscape phenology from unmanned aerial vehicle (UAV) photography *Agric. Forest Meteorol.* **248** 397–407
- [117] Berra E F, Gaulton R and Barr S 2019 Assessing spring phenology of a temperate woodland: A multiscale comparison of ground, unmanned aerial vehicle and Landsat satellite observations *Remote Sens. Environ.* **223** 229–42
- [118] D’Odorico P, Besik A, Wong C Y S, Isabel N and Ensminger I 2020 High-throughput drone-based remote sensing reliably tracks phenology in thousands of conifer seedlings *New Phytol.* **226** 1667–81
- [119] Prev y J et al 2017 Greater temperature sensitivity of plant phenology at colder sites: implications for convergence across northern latitudes *Glob. Change Biol.* **23** 2660–71
- [120] Fawcett D and Anderson K 2019 Investigating impacts of calibration methodology and irradiance variations on lightweight drone-based sensor derived surface reflectance products. in: remote sensing for agriculture, ecosystems, and hydrology XXI International Society for Optics and Photonics (available at: (<https://www.spiedigitallibrary.org/conference-proceedings-of-spie/11149/111490D/Investigating-impacts-of-calibration-methodology-and-irradiance-variations-on-lightweight/10.1117/12.2533106.short>) (Accessed 11 July 2020) pp 111490D
- [121] Huang S, Tang L, Hupy J P, Wang Y and Shao G 2020 A commentary review on the use of normalized difference vegetation index (NDVI) in the era of popular remote sensing *J. Forestry Res.* (<https://doi.org/10.1007/s11676-020-01155-1>)
- [122] Zhang Q, Yao T, Huemmrich K F, Middleton E M, Lyapustin A and Wang Y 2020 Evaluating impacts of snow, surface water, soil and vegetation on empirical vegetation and snow indices for the Utqiagvik tundra ecosystem in Alaska with the LVS3 model *Remote Sens. Environ.* **240** 111677
- [123] Aasen H, Honkavaara E, Lucieer A and Zarco-Tejada P J 2018 Quantitative remote sensing at ultra-high resolution with UAV Spectroscopy: a review of sensor technology, measurement procedures, and data correction workflows *Remote Sens.* **10** 1091
- [124] Strahler A H, Woodcock C E and Smith J A 1986 On the nature of models in remote sensing *Remote Sens. Environ.* **20** 121–39
- [125] Chen J M 1999 Spatial scaling of a remotely sensed surface parameter by contexture *Remote Sens. Environ.* **69** 30–42
- [126] Chen J M, Chen X and Ju W 2013 Effects of vegetation heterogeneity and surface topography on spatial scaling of net primary productivity *Biogeosciences* **10** 4879–96
- [127] Garrigues S, Allard D, Baret F and Weiss M 2006 Influence of landscape spatial heterogeneity on the non-linear estimation of leaf area index from moderate spatial resolution remote sensing data *Remote Sens. Environ.* **105** 286–98